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(54) Oil resistant silicone sealants

(57) The invention relates to a RTV silicone composition having increased hot oil resistance upon curing. The RTV silicone composition comprises a polydiorganosiloxane, an acyloxy-functional crosslinker, a metal salt of a carboxylic acid catalyst, a filler, and a metal acetylacetonate. In a preferred embodiment a metal oxide is also added.

Description

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The use of room temperature vulcanizing (RTV) silicone sealants for creating formed-in-place gaskets is well known in both original equipment manufacture and in repair and maintenance. A problem with certain conventional silicone sealants is their tendency in the presence of hot oil to lose structural integrity leading to seal failure. Therefore, it is desirable to have silicone sealants providing increased oil resistance as demonstrated by increased tensile strength in the presence of hot oil.

The prior art is represented by U.S. Patent No. 4,123,472; U.S. Patent No. 4,680,363; U.S. Patent No. 4,833,037; U.S. Patent No. 4,257,932 and U.S. Patent No. 4,797,462.

The present inventors have unexpectedly determined that the addition of a metal acetylacetonate, preferably along with a metal oxide, to a RTV silicone composition comprising a polydiorganosiloxane, acyloxy-functional crosslinker, filler, and tin catalyst results in a cured composition having increased hot oil resistance. In addition, these compositions may find use when cured as sealants, adhesives, gaskets, coatings, molding and potting compounds and gels.

The RTV silicone composition which is curable in the presence of moisture comprises a polydiorganosiloxane, an acyloxy-functional crosslinker, a metal salt of a carboxylic acid catalyst, a filler, and a metal acetylacetonate.

This invention is a RTV silicone composition which is curable in the presence of moisture and has increased hot oil resistance upon curing comprising:

- (A) 100 parts by weight of a polydiorganosiloxane in which the terminal groups are selected from silanol and triorganosilyl groups, provided at least 60 mole percent of the terminal groups are silanol groups;
- (B) 1 to 15 parts by weight of an acyloxy-functional crosslinking agent described by the formula $R_mSi(OCOR')_{4-m}$ where R is a monovalent hydrocarbon radical selected from 1 to 12 carbon atoms, each R' is an independently selected monovalent hydrocarbon radical from 1 to 12 carbon atoms and m is 0 or 1;
- (C) 0.001 to 1 part by weight of a metal salt of a carboxylic acid catalyst;
- (D) 5 to 150 parts by weight of a filler; and
- (E) an effective amount of a metal acetylacetonate.

Component (A) is a polydiorganosiloxane having terminal groups selected from silanol and triorganosilyl groups, provided at least 60 mole percent of the terminal groups are silanol groups. The polydiorganosiloxane may be one type of polymer or a mixture of different polymers. The polydiorganosiloxanes may have a linear structure and may be homopolymers or copolymers. In addition, the organic groups linked to any particular silicon atom may be the same or different. The organic groups of the polydiorganosiloxane can include any monovalent hydrocarbon group or any monovalent halohydrocarbon group comprising 1 to 18 carbon atoms. Preferred organic groups are methyl, ethyl, propyl, phenyl, vinyl and 3,3,3-trifluoropropyl, with methyl being most preferred.

The terminal groups of the polydiorganosiloxane are selected from silanol and triorganosilyl groups. The organic substituent of the triorganosilyl groups can include any monovalent hydrocarbon group or any monovalent halohydrocarbon group comprising 1 to 18 carbon atoms. The preferred organic substituent of the triorganosilyl groups are methyl, ethyl, propyl, phenyl, vinyl, and 3,3,3-trifluoropropyl, with methyl being most preferred.

Although the terminal groups are selected from silanol and triorganosilyl groups, it is required that at least 60 mole percent of the terminal groups be silanol groups in order to obtain the desired properties of the composition when cured. Preferably, 80 to 100 mole percent of the terminal groups of the polydiorganosiloxane are silanol groups.

The viscosity of the polydiorganosiloxane may be from 1 to 150 Pa •s at 25°C. The preferred viscosity of the polydiorganosiloxane is from 5 to 100 Pa •s at 25°C.

The methods of manufacture of the silanol terminated polydiorganosiloxanes are well known in the art. One common method is based upon the hydrolysis of diorganodichlorosilane, the separation of a diorganotetrasiloxane cyclic material from the hydrolysis mixture, and the subsequent polymerization of the cyclic material to the polydiorganosiloxane through the use of an alkaline catalyst. The triorganosilyl terminated polydiorganosiloxanes are also prepared by known methods described in U.S. Patent No. 3,274,145.

Component (B) is an acyloxy-functional crosslinking agent described by the formula $R_mSi(OCOR')_{4-m}$ where R is a monovalent hydrocarbon radical selected from 1 to 12 carbon atoms, each R' is an independently selected monovalent hydrocarbon radical from 1 to 12 carbon atoms and m is 0 or 1.

The monovalent hydrocarbon radicals represented by R may be linear or branched and can include alkyl radicals such as methyl, ethyl, isopropyl or hexyl; alkenyl radicals such as vinyl, allyl or hexenyl; alkynal radicals such as propargyl; cycloaliphatic radicals such as cyclopentyl, cyclohexyl or cyclohexenyl; aromatic radicals such as phenyl or tolyl; and aralkyl radicals such as benzyl or beta-phenylethyl. R is preferably methyl or ethyl.

Each R' is an independently selected monovalent hydrocarbon radical from 1 to 12 carbon atoms. Examples of R' include those provided for R. R' is preferably methyl or ethyl, with methyl being most preferred.

The acyloxy-functional crosslinker may have trifunctionality, as occurs when m is 1 or tetrafunctionality, as occurs

when m is 0. It is preferred that m be 1.

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Examples of preferred acyloxy-functional crosslinkers include methyltriacetoxysilane, ethyltriacetoxysilane, methyltris-benzoxysilane, vinyltriacetoxysilane and methyl-tris-2-ethylhexanoxy silane.

The acyloxy-functional crosslinker can be present in the present composition in amounts from 1 to 15 parts by weight based on 100 parts by weight of the polydiorganosiloxane, and preferably from 2 to 10 parts by weight on the same basis. The acyloxy-functional crosslinker may be added as a single species or as a mixture of two or more species.

Component (C) is a metal salt of a carboxylic acid catalyst. The metal component of the catalyst is preferably tin, but may be lead, chromium, antimony, iron, cadmium, barium, titanium, bismuth or magnesium. Examples of suitable metal salts of carboxylic acids include tin naphthenate, lead octoate, tin octoate, iron stearate, tin oleate, dibutyltindilaurate, dibutyltindiacetate, dibutyl tin oxide, and dimethyl tin bis-neodecanoate. Preferred catalysts are dibutyltindilaurate, dibutyltindiacetate, and dimethyl tin bis-neodecanoate. The most preferred catalysts are dibutyltindilaurate and dibutyltindiacetate.

The catalyst can be present in the present composition in amounts from 0.001 to 1 part by weight based on 100 parts by weight of the polydiorganosiloxane, and preferably from 0.01 to 0.3 part by weight on the same basis. It is most preferred to have 0.01 to 0.2 part by weight catalyst present in the composition on the same basis. The catalyst may be added as a single species or as a mixture of two or more species.

In order to obtain the desired physical properties of the present composition when cured, a filler (Component (D)) is added to the composition. This filler may include reinforcing fillers, semi-reinforcing fillers, or non-reinforcing fillers also called extending fillers, or any combination thereof. One or more of each type of filler is also acceptable.

Reinforcing fillers include any finely divided form of silica which can be prepared by precipitation or a pyrogenic process and may be treated or untreated. Generally, methods of treating the silica can include mixing the silica with polycyclosiloxane such as disclosed in U.S. Patent No. 2,938,009 or U.S. Patent No. 3,004,859 or silazanes as disclosed in U.S. Patent 3,635,743. The treating agents may also include liquid hydroxyl-terminated polydiorganosiloxanes which can contain an average of about 2 to 20 repeating units and can contain at least one alkenyl unit. The treating agent may also be an alkylhalosilane, such as dimethyldichlorosilane or an alkoxysilane. Carbon black is also useful as a reinforcing filler in this invention. Semi-reinforcing fillers include barium sulfate and crystalline silica. Non-reinforcing or extending fillers include glass spheres, wollastonite, diatomaceous earths, clay and talc. The preferred fillers are untreated fumed silica and treated fumed silica.

The filler can be present in the present composition in amounts from 5 to 150 parts by weight based on 100 parts by weight of the polydiorganosiloxane, and preferably, from 5 to 30 parts by weight on the same basis. The filler may be added as a single species or as a mixture of two or more species.

Component (E) is a metal acetylacetonate. The inventors have unexpectedly determined that the addition of a metal acetylacetonate to an RTV silicone composition provides the composition upon curing with increased hot oil resistance.

The metal portion of the metal acetylacetonate is preferably copper, iron or zirconium, with copper being most preferable, but may be selected from copper, chromium, iron, aluminum, zinc, titanium and zirconium. Examples of Component (E) include copper II acetylacetonate, ferric acetylacetonate, chromium III acetylacetonate and zinc acetylacetonate.

An effective amount of the metal acetylacetonate is an amount which provides increased hot oil resistance to the RTV silicone composition upon curing. Typically, 0.01 to 5 parts by weight metal acetylacetonate based on 100 parts by weight of the polydiorganosiloxane is effective. It is preferable to add 0.05 to 1.5 parts by weight metal acetylacetonate on the same basis, with 0.1 to 0.6 part by weight metal acetylacetonate on the same basis being most preferred. The metal acetylacetonate may be added as a single species or as a mixture of two or more species.

In a preferred embodiment Component (F), a metal oxide, is also added to the RTV silicone composition. The metal portion of the metal oxide is most preferably calcium, but may also be selected from calcium, zinc, magnesium, aluminum, zirconium, barium, titanium, strontium and copper. Examples of Component (F) include calcium oxide, zinc oxide, magnesium oxide, aluminum oxide, zirconium oxide, barium oxide, titanium IV dioxide and copper II oxide.

The metal oxide can be present in the present composition in amounts from 0.01 to 10 parts by weight based on 100 parts by weight of the polydiorganosiloxane. It is preferable to add from 0.01 to 1.5 parts by weight on the same basis and, most preferably, from 0.01 to 1 part by weight on the same basis. The metal oxide may be added as a single species or as a mixture of two or more species.

In preferred embodiments, the present compositions can also optionally include up to 2 parts by weight of an adhesion promoter based on 100 parts by weight of the polydiorganosiloxane. Examples of useful adhesion promoters include ethylpolysilicate and glycidoxypropyltrimethoxysilane.

In addition to the above ingredients, the present composition may include additives which impart or enhance certain properties of the cured composition or facilitate processing of the curable composition. Typical additives include plasticizers, functional and non-functional diluents, pigments, dyes, heat and/or ultraviolet light stabilizers, flame-retard-

ant agents, thickeners, rheology modifiers, and preservatives. The effect of any such additives should be evaluated as to their result and impact on other properties of the composition.

The RTV silicone composition of this invention may be prepared by mixing all the ingredients together in amounts as specified to provide a composition which is stable in the absence of moisture and which cures to an elastomeric material when exposed to moisture.

These compositions are preferably prepared by mixing the ingredients under anhydrous conditions. This means that the amounts of moisture or water in the ingredients used should be minimized and the conditions of mixing should minimize the amounts of moisture or water allowed into the system. Excess moisture or water may have deleterious effects on the composition, such as causing curing in the storage package or reduced cured properties.

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The present compositions may be prepared as either a one package system or a two (or multi) package system. With a one package system, all the ingredients are mixed together and stored in an anhydrous state. With a two or multi package system the ingredients may be stored separately and then mixed prior to use. It is often convenient to form a "base" blend comprising a polydiorganosiloxane, a filler and, if needed, a plasticizer (I). Another blend comprising curing agents such as the crosslinker and catalyst may also be mixed together (II) and still another blend comprising additional ingredients including Components (E) and (F) may also be mixed together (III). As described above, the components comprising (I), (II) and (III) may be mixed together or separately. If mixed and stored together a one part sealant results. Obviously a two or multi package system is possible if (I), (II), and (III) are mixed and packaged separately. It is also possible to mix and store two of the components together, i.e., (I) and (III) and later mix in (III). In a preferred embodiment of the present invention, it is preferred to mix and store (I) and (III) together and then later mix in (III) as required.

The present RTV silicone compositions may be used as gasket materials for sealing gaps or seams in a given substrate or for sealing between substrates. Due to the oil resistance of the present RTV silicone compositions when cured these compositions are preferably used for sealing gaps or substrates having surfaces exposed to oil or hot oil. However, these RTV silicone compositions may also be used to seal gaps or substrates which may not have surfaces exposed to oil or hot oil.

A method for sealing using the present composition comprises (i) providing at least two substrates having a plurality of surfaces; (ii) applying a layer of a RTV silicone composition which is curable in the presence of moisture, to at least a portion of at least one of the surfaces of at least one of the substrates, the RTV silicone composition comprising Components (A)-(E) and preferably also Component (F), as described above; (iii) bringing at least two of the substrates, where at least one of the substrates has the RTV silicone composition applied thereto, into proximity so as to form a gasket of the RTV silicone composition therebetween; and (iv) exposing the gasket of step (iii) to moisture to effect curing of the RTV silicone composition.

The substrates useful for this preferred embodiment can be metal materials, composites or certain plastic materials. Preferably the substrates are metal materials.

The following examples are presented for illustrative purposes and should not be construed as limiting the invention which is delineated in the claims.

In the following examples, unless otherwise noted, the compositions were mixed at 23°C±2°C and 50%±5% relative humidity; cast, and then cured for 7 days at 23°C±2°C and 50%±5% relative humidity before oil immersion testing. The cured samples were then immersed in oil for 14 days at 150°C per ASTM D471 except the samples were cooled to 23°C±2°C in the test fluid rather than in a cooling fluid. The oil used for immersion testing was Goodwrench 5W30 SAE SH/SG. The durometer (Shore A) results provided in each table were obtained using the method described in ASTM D2240 with the following exceptions: 2.54 cm x 1.91 cm tabs having a thickness of 0.64 cm were tested and an average of 5 readings was reported. The tensile, 100% modulus, and elongation results provided in each table were obtained using the method described in ASTM D412 with each tensile, 100% modulus, and elongation value reported as the average of 3 samples. Adhesion testing was conducted according to ASTM D903 with the following exceptions: 1) the speed of testing was 5.08 cm/min., 2) substrates were cleaned with MIBK (methylisobutylketone) followed by an acetone rinse and 3) the flexible substrate was 1.27 cm wide. Volume swell % results were obtained using the method described in ASTM 471. Also, as used herein "part" or "parts" are by weight and "AcAc" refers to acetylacetonate.

Prior to use, the metal oxide powder was dried at 105°C. for a minimum of 2 hours. All metal oxides used in the examples with the exception of copper II oxide were obtained from Fisher Scientific (Pittsburgh, PA). Copper II oxide was obtained from Aldrich Chemical (Milwaukee, WI). Copper II AcAc was obtained from Acros Organics (Pittsburgh, PA) and Aldrich Chemical. Chromium III AcAc, Iron III AcAc and Aluminum AcAc were obtained from Acros Organics. Zinc AcAc Hydrate, Titanium IV Oxide AcAc and Zirconium AcAc were obtained from Aldrich Chemical. The fumed silica was obtained from Cabot Corp (Tuscola, IL). The dibutyl tin diacetate was obtained from Atochem Corporation (Philadelphia, PA). The silane and siloxane compounds were obtained from Dow Corning Corporation (Midland, MI). Ethylpolysilicate was obtained from Huls of America, Inc. (Piscataway, NJ).

Example 1

Formulations were prepared as shown in Table 1.1. Ingredients numbered as 1, 2, 3 and 8 were combined with a compounder at approximately 70°C., packaged into cartridges and then deaired. Ingredients numbered 4, 5, 6, 7 and 9 were mixed together and then injected into and mixed with the mixture of ingredients 1, 2, 3 and 8 using a Semco mixer (Model 388, Products Research and Chemical Corp, Glendale, CA). Samples were cast, cured and then tested in hot oil. Formulations 1A and 1C are controls while Formulations 1B and 1D have added copper acetylacetonate and calcium oxide. The post oil immersion tensile strength improved from less than 0.09 N/mm² without the additives to greater than 0.75 N/mm² with the additives. The results are provided in Tables 1.2(a) and 1.2(b).

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Table 1.1

Formulations

Form. 1A*

100.00

13.86

6.19

4.80

2.40

0.00

0.00

0.03

0.64

Form. 1B

100.00

13.79

6.19

4.80

2.4

0.24

0.36

0.03

0.64

Form. 1C*

100.00

12.38

5.55

4.72

2.36

0.00

0.00

0.03

0.63

Form. 1D

100.00

11.69

5.23

4.68

2.34

0.24

0.35

0.03

0.63

15

20

25

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6.

7.

* Formulations 1A and 1C are the controls, with no additives included.

17.5 Pa • s, 85% silanol terminated and 15% trimethyl

Untreated fumed silica having a surface area of approx-

Mixture of methyl(~45%) and ethyl(~55%) triacetoxysi-

0.1 Pa • s Trimethyl Endblocked Polydimethylsiloxane

Approximately 0.075 Pa • s hydroxy-terminated poly-

Component (parts)

imately 200 m²/g

dimethylsiloxane

Calcium Oxide

Dibutyl tin diacetate

Blue Silicone Paste**

Copper acetylacetonate

lane

terminated polydimethylsiloxane

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Table 1:2(a)

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Initial Results Prior to Oil Immersion										
Form. 1A* Form. 1B Form. 1C* Form 1D										
Durometer, Shore A	23	28	28	23						
Tensile Strength, N/mm ²	1.48	1.69	2.16	1.48						
Elongation, %	349	366	460	349						
Modulus @ 100%, N/mm ²	0.50	0.54	0.56	0.50						

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* Formulations 1A and 1C are the controls, with no additives included.

^{**} Ferro SV1232, Ferro Corp. (Edison, NJ)

Table 1.2(b)

5W30 Oil Immersion Results Form. 1C* Form. 1A* Form. 1B Form 1D Durometer, Shore A 0 0 6 4 Tensile Strength, N/mm² 0.09 0.75 0.04 0.71 Elongation, % 442 26 233 502 Modulus @ 100%, N/mm² N/A 0.28 0.21 80.0 Volume Swell % 55 51 35 52

Example 2

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The ingredients listed in Table 2.1 were mixed and tested as described in Example 1. Ingredients numbered as 1, 2, 3 and 7 were combined with a compounder at approximately 70°C. The mixture was then pigmented by mixing in ingredient 8, packaged into cartridges and then deaired. Ingredients numbered 4, 5 and 6 were mixed together and then injected into and mixed with the mixture of ingredients 1, 2, 3, 7 and 8 using a Semco mixer. The post oil immersion tensile strength was improved with the addition of the copper AcAc or calcium oxide added individually, however, the combined effect is greater than either with copper AcAc or calcium oxide added individually. The results are provided in Tables 2.2(a) and 2.2(b). 25

Table 2.1

1.17.5 Pa • s, 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane100.00100.002.Untreated fumed silica having a surface area of approximately 200 m²/g13.3613.363.Mixture of methyl(~45%) and ethyl(~55%) triacetoxysilane5.975.974.12 Pa • s Trimethyl Endblocked Polydimethylsiloxane1.981.985.Copper acetylacetonate0.000.310.006.Calcium Oxide0.000.000.62				•	
•	Component (parts)	Form. 2A*	Form. 2B	Form. 2C	Form. 2D
1.		100.00	100.00	100.00	100.00
2.		13.36	13.36	13.36	13.86
3.		5.97	5.97	5.97	6.19
4.	12 Pa · s Trimethyl Endblocked Polydimethylsiloxane	1.98	1.98	1.98	2.05
5.	Copper acetylacetonate	0.00	0.31	0.00	0.64
6.	Calcium Oxide	0.00	0.00	0.62	0.64
7.	Dibutyl tin diacetate	0.02	0.02	0.02	0.03
8.	Black Pigment**	2.48	2.48	2.48	2.57

^{*} Formulation 2A is the control, with no additives included.

^{*} Formulations 1A and 1C are the controls, with no additives included.

^{**} Paste of LV1011 Lampblack Powder, Harcross Pigments (Fairview Heights, IL)

Table 2.2(a)

Initial Results Prior to Oil Immersion							
	Form. 2A*	Form. 2B	Form. 2C	Form. 2D			
Durometer, Shore A	25	26	26	27			
Tensile Strength, N/mm ²	2.02	1.73	2.01	1.70			
Elongation, %	473	426	469	405			
Modulus @ 100%, N/mm ²	0.47	0.49	0.48	0.50			

^{*} Formulation 2A is the control, with no additives included.

Table 2.2(b)

5W30 Oil Immersion Results									
Form. 2A* Form. 2B Form. 2C Form. 2D									
Durometer, Shore A	0	0	0	6					
Tensile Strength, N/mm ²	0.00	0.28	0.42	1.05					
Elongation, %	. 0	849	589	574					
Modulus @ 100%, N/mm ²	0.00	0.08	0.16	0.26					
Volume Swell %	N/A	57	58	54					

^{*} Formulation 2A is the control, with no additives included.

Example 3

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The ingredients listed in Table 3.1 were mixed and tested as described in Example 1. Ingredients numbered as 1, 2, 3 and 7 were combined with a compounder at approximately 70°C., packaged into cartridges and then deaired. Ingredients numbered 4, 5 and 6 were mixed together and then injected into and mixed with the mixture of ingredients 1, 2, 3 and 7 using a Semco mixer. In this example, different amounts of the copper AcAc and calcium oxide were used individually (Tables 3.1(a) and 3.1(b) respectively) and in combination (Table 3.1(c)). The results are provided in Tables 3.2(a) and 3.2(b).

Table 3.1(a)

		1000.0	()				
		Addition of Copper Ace	tylacetonat	e Only			<u>.</u> . <u>.</u>
	_	Component (parts)	3A	3B	3C	3D	3E
45 ,	1.	17.5 Pa • s, 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane	100.0 0	100.0 0	100.0 0	100.0 0	100.0 0
	2.	Untreated fumed silica having a surface area of approximately 200 m ² /g	22.21	22.21	22.21	22.21	22.21
50	3.	Mixture of methyl(~45%) and ethyl(~55%) triacetox-ysilane	9.15	9.15	9.15	9.15	9.15
	4.	12 Pa · s Trimethyl Endblocked Polydimethylsiloxane	1.97	1.97	1.97	1.97	1.97
	5.	Copper acetylacetonate	0.32	0.52	0.99	1.31	1.97
<i>55</i>	6.	Calcium Oxide	0.00	0.00	0.00	0.00	0.00
	7.	Dibutyl tin diacetate	0.05	0.05	0.05	0.05	0.05

Table 3.1(b)

	Addition of Calcium Oxide Only							
	Component (parts)	3F	3G	3H	31			
1.	17.5 Pa • s, 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane	100.0 0	100.0 0	100.0 0	100.0 0			
2.	Untreated fumed silica having a surface area of approximately 200 m ² /g	22.21	22.21	22.21	22.21			
3.	Mixture of methyl(~45%) and ethyl(~55%) triacetoxysilane	9.15	9.15	9.15	9.15			
4.	12 Pa · s Trimethyl Endblocked Polydimethylsiloxane	2.10	2.10	2.10	2.10			
5.	Copper acetylacetonate	0.00	0.00	0.00	0.00			
6.	Calcium Oxide	0.33	0.65	0.98	1.32			
7.	Dibutyl tin diacetate	0.05	0.05	0.05	0.05			

Table 3.1(c)

	iable 0.1(0)							
	Addition of Copper Acetylacetonate and Calcium Oxide							
	Component (parts)	3J	ЗК					
1.	17.5 Pa • s, 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane	100.00	100.00					
2.	Untreated fumed silica having a surface area of approximately 200 m ² /g	22.21	22.21					
3.	Mixture of methyl(~45%) and ethyl(~55%) triacetoxysilane	9.15	9.15					
4.	12 Pa · s Trimethyl Endblocked Polydimethylsiloxane	2.10	2.63					
5.	Copper acetylacetonate	0.33	0.66					
6.	Calcium Oxide	0.98	0.66					
7.	Dibutyl tin diacetate	0.05	0.05					

Table 3.2(a)

Initial Results Prior to Oil Immersion								
Formulation	Durometer Shore A	Tensile N/mm ²	Elongation %	100% Modulus N/mm ²				
3A	46	2.47	240	1.30				
3B	46	2.70	269	1.27				
3C	48	2.79	280	1.27				
3D	49	2.42	239	1.28				
3E	49	2.43	244	1.28				
3F	45	2.73	306	1.14				

Table 3.2(a) (continued)

Initial Results Prior to Oil Immersion							
Formulation	Durometer Shore A	Tensile N/mm ²	Elongation %	100% Modulus N/mm ²			
3G	46	2.68	299	1.14			
3H	44	2.83	318	1.13			
31	43	2.97	328	1.12			
3.J	44	2.73	283	1.21			
3K	47	2.47	251	1.27			

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Table 3.2(b)

Elongation %

354

294

329

311

305

260

271

317

266

290

282

100% Modulus

· N/mm²

0.42

0.49

0.53

0.52

0.5

0.45

0.61

0.59

0.59

0.68

0.74

Volume Swell %

36

35

35

35

35

37

37

37

34

32

29

5W30 Oil Immersion Results 20 Formulation Durometer Shore A Tensile N/mm² 13 **3A** 0.88 3B 15 0.92 25 3C 17 1.14 15 3D 1.05 14 3E 0.96 30 3F 14 0.72 17 3G 1.07 3H 18 1.32 35 31 19 1.27

22

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Example 4

3J

3K

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The ingredients listed in Table 4.1 were mixed and tested as described in Example 1. Ingredients numbered as 1, 2, 3 and 8 were combined with a compounder at approximately 70°C., packaged into cartridges and then deaired. Ingredients numbered 4, 5, 6, 7, 9 and 10 were mixed together and then injected into and mixed with the mixture of ingredients 1, 2, 3 and 8 using a Semco mixer. In this example, an adhesion promoter (10) was also added. The adhesion promoter did not result in degraded oil resistance. The results are provided in Tables 4.2(a) and 4.2(b).

1.45

1.50

Table 4.1

	Formulations				-
	Component (parts)	Form. 4A	Form. 4B	Form. 4C	Form. 40
1.	17.5 Pa • s, 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane	100.00	100.00	100.00	100.00
2.	Untreated fumed silica having a surface area of approximately 200 m ² /g	13.89	13.93	15.05	15.15
3.	Mixture of methyl(~45%) and ethyl(~55%) triacetoxysilane	6.20	6.22	6.78	6.82
4.	0.1 Pa · s Trimethyl Endblocked Polydimethylsiloxane	4.81	4.83	2.09	2.11
5.	Approximately 0.075 Pa • s hydroxy-terminated poly- dimethylsiloxane	2.41	2.41	2.45	2.46
6.	Copper acetylacetonate	0.24	0.25	0.25	0.25
7.	Calcium Oxide	0.36	0.36	0.37	0.37
8.	Dibutyl tin diacetate	0.03	0.03	0.03	0.03
9.	Blue Silicone Paste*	0.48	0.48	0.48	0.49
10.	Ethylpolysilicate	0.64	1.29	0.65	1.32

^{*} Ferro SV1232, Ferro Corp. (Edison, NJ)

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Table 4.2(a)

Initial Results Prior to Oil Immersion								
	Form. 4A	Form. 4B	Form. 4C	Form. 4D				
Durometer, Shore A	25	25	28	28				
Tensile Strength, N/mm ²	1.70	1.95	2.01	2.13				
Elongation, %	425	452	429	469				
Modulus @ 100%, N/mm ²	0.48	0.44	0.52	0.52				
Adhesion to 1010 steel/aluminum, pli (N/mm)	6/28 (1.05/4.9)	14/32 (2.45/5.6)	6/34 (1.05/5.95)	22/38 (3.85/6.65)				

Table 4.2(b)

5W30 Oil Immersion Results						
	Form. 4A	Form. 4B	Form. 4C	Form. 4D		
Durometer, Shore A	3	4	5	6		
Tensile Strength, N/mm ²	0.70	0.74	0.91	0.97		
Elongation, %	572	557	560	583		
Modulus @ 100%, N/mm ²	0.19	0.21	0.23	0.23		
Volume Swell %	55	52	49	48		

Example 5

The ingredients listed in Table 5.1 were mixed and tested as described in Example 1. Ingredients numbered as 1, 2, 3 and 8 were combined with a compounder at approximately 70°C., packaged into cartridges and then deaired. Ingredients numbered 4, 5, 6, 7, 9 and were mixed together and then injected into and mixed with the mixture of ingredients 1, 2, 3 and 8 using a Semco mixer. In this 5.2(a)(1) and (b)(1). Each Form. 5B incorporates calcium oxide with a metal AcAc as described in the results Tables 5.2(a)(2) and (b)(2).

Table 5.1

	Formulations		
	Components (parts)	Form. 5A	Form 5B
1	17.5 Pa • s viscosity 85% silanol terminated and 15% trimethyl terminated polydimethylsiloxane	100	100 ,
2	Untreated fumed silica having a surface area of approximately 200 m ² /g	13.09	13.09
3	Mixture of methyl(~45%) and ethyl(~55%) triacetoxysilane	5.86	5.86
4	0.1 Pa s Trimethyl Endblocked Polydimethylsiloxane	4.76	4.76
5	Approximately 0.075 Pa • s viscosity hydroxy-terminated polydimethylsiloxane.	2.38	2.38
6a	copper acetylacetonate	0.24	0.0
6b	metal acetylacetonate (see Tables 5.2(a)(1) and (b)(1))	0.0	0.24
7a	calcium oxide	0.0	0.36
7b	metal oxide (see Tables 5.2(a)(2) and (b)(2))	0.36	0.0
8	Dibutyl tin diacetate	0.03	0.03
9	Blue Silicone Paste*	0.44	0.44
10	ethylpolysilicate	0.59	0.59

^{*} Ferro SV1232, Ferro Corp. (Edison, NJ)

Table 5.2(a)

	Initial Results (1) Samples containing a	Prior to Oil Immer a Metal AcAc and (
Additive	Durometer (Shore A)	Tensile N/mm ²	Elongation%	100% Modulus N/mm ²
Chromium III AcAc	24.0	1.89	392	0.48
Iron III AcAc	28.0	1.64	348	0.56
Aluminum AcAc	29.0	1.44	333	0.55
Zirconium AcAc	30.0	1.59	289	0.63
Zinc AcAc Hydrate	26.0	1.53	347	0.52
Titanium IV Oxide AcAc	33.0	1.59	270	0.68
(2) 5	Samples containing Copp	er AcAc and a Me	tal Oxide (Form.(A))
Additive	Durometer (Shore A)	Tensile N/mm ²	Elongation%	100% Modulus N/mm ²
Zinc Oxide	26.0	1.65	393	0.50
Magnesium Oxide	26.0	1.56	400	0.48
Titanium Dioxide	26.0	2.04	484	0.48
Barium Oxide	24.0	1.64	435	0.45
Aluminum Oxide	24.0	1.48	398	0.46
Zirconium Oxide	25.0	1.65	409	0.48
Copper II Oxide	30.0	1.88	392	0.54

100% Modulus

100% Modulus

0.13

Volume Swell %

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Volume Swell %

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		Table 5.2(b)						
5		5W30 Oil Immersion Results (1) Samples containing a Metal AcAc and CaO (Form.(B))						
		Additive	Durometer (Shore A)	Tensile N/mm ²	Elongation%	100% Modulus N/mm ²		
	10	Chromium III AcAc	0	0.60	829	0.12		
•		Iron III AcAc	. 3	0.92	539	0.23		
		Aluminum AcAc	0	0.40	705	0.11		
15	15	Zirconium AcAc	. 5	0.81	382	0.24		
	ıs	Zinc AcAc Hydrate	4	0.72	583	0.19		
		Titanium IV Oxide AcAc	. 4	0.76	446	0.23		
•	20	(2) Samples containing Copper AcAc and a Metal Oxide (Form.(A))						
		Additive	Durometer (Shore A)	Tensile N/mm ²	Elongation%	100% Modulus N/mm ²		
	25	Zinc Oxide	0	0.46	789	0.12		
25	Magnesium Oxide	4	0.78	632	0.18			
		Titanium Dioxide	1	0.59	807	0.12		
		Barium Oxide	. 0	0.54	715	0.12		
	30	Aluminum Oxide	0	0.43	610	0.12		
		Zirconium Oxide	1	0.51	706	0.12		

Claims

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Copper II Oxide

A RTV silicone composition which is curable in the presence of moisture, comprising:

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(A) 100 parts by weight of a polydiorganosiloxane in which the terminal groups are selected from silanol and triorganosilyl groups, provided at least 60 mole percent of the terminal groups are silanol groups;

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- (B) 1 to 15 parts by weight of an acyloxy-functional crosslinking agent described by the formula R_mSi(OCOR')₄₋ m where R is a monovalent hydrocarbon radical selected from 1 to 12 carbon atoms, each R' is an independently selected monovalent hydrocarbon radical from 1 to 12 carbon atoms and m is 0 or 1;
- (C) 0.001 to 1 part by weight of a metal salt of a carboxylic acid catalyst;
- (D) 5 to 150 parts by weight of a filler; and

- (E) an effective amount of a metal acetylacetonate.
- The RTV silicone composition of claim 1 comprising 0.01 to 5 parts by weight of the metal acetylacetonate based on 100 parts by weight of the polydiorganosiloxane, where the metal portion of the metal acetylacetonate is 50 selected from copper, chromium, iron, aluminum, zinc, titanium and zirconium.
 - The RTV silicone composition of claim 1 where the polydiorganosiloxane is a polydimethylsiloxane in which 80 to 100 mole percent of the terminal groups are silanol groups; R and R' of the acyloxy-functional crosslinking agent. are independently selected from methyl and ethyl, and m is 1; the catalyst is a tin salt of a carboxylic acid and the filler is a silica filler.
 - The RTV silicone composition of claim 1 further comprising (F) 0.01 to 10 parts by weight of a metal oxide, based

on 100 parts by weight of the polydiorganosiloxane.

- 5. The RTV silicone composition of claim 4 comprising 0.01 to 1.5 parts by weight of the metal oxide based on 100 parts by weight of the polydiorganosiloxane, where the metal portion of the metal oxide is selected from the group consisting of calcium, zinc, magnesium, aluminum, zirconium, barium, titanium, strontium and copper.
- **6.** A method of sealing, comprising the steps of:

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- (i) providing at least two substrates having a plurality of surfaces;
- (ii) applying a layer of a RTV silicone composition which is curable in the presence of moisture, to at least a portion of at least one of the surfaces of at least one of the substrates, the RTV silicone composition comprising:
 - (A) 100 parts by weight of a polydiorganosiloxane in which the terminal groups are selected from silanol and triorganosilyl groups, provided at least 60 mole percent of the terminal groups are silanol groups,
 - (B) 1 to 15 parts by weight of an acyloxy-functional crosslinking agent described by the formula $R_mSi(OCOR')_{4-m}$ where R is a monovalent hydrocarbon radical selected from 1 to 12 carbon atoms, each R' is an independently selected monovalent hydrocarbon radical from 1 to 12 carbon atoms and m is 0 or 1,
 - (C) 0.001 to 1 part by weight of a metal salt of a carboxylic acid catalyst,
 - (D) 5 to 150 parts by weight of a filler, and
 - (E) an effective amount of a metal acetyl-acetonate,
- (iii) bringing at least two of the substrates, where at least one of the substrates has the RTV silicone composition applied thereto, into proximity so as to form a gasket of the RTV silicone composition therebetween; and (iv) exposing the gasket of step (iii) to moisture to effect curing of the RTV silicone composition.
- 7. The method of claim 6 where the RTV silicone composition comprises about 0.01 to 5 parts by weight of the metal acetylacetonate based on 100 parts by weight of the polydiorganosiloxane, where the metal portion of the metal acetylacetonate is selected from copper, chromium, iron, aluminum, zinc, titanium and zirconium.
- 8. The method of claim 6 where in the RTV silicone composition, the polydiorganosiloxane is a polydimethylsiloxane in which 80 to 100 mole percent of the terminal groups are silanol groups; R and R' of the acyloxy-functional crosslinking agent are independently selected from methyl and ethyl and m is 1; the catalyst is a tin salt of a carboxylic acid; and the filler is a silica filler.
- 9. The method of claim 6 where the RTV silicone composition further comprises (F) 0.01 to 10 parts by weight of a metal oxide, based on 100 parts by weight of the polydiorganosiloxane.

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(12)

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(54) Oil resistant silicone sealants

(57) The invention relates to a RTV silicone composition having increased hot oil resistance upon curing. The RTV silicone composition comprises a polydiorganosiloxane, an acyloxy-functional crosslinker, a metal salt of a carboxylic acid catalyst, a filler, and a metal acetylacetonate. In a preferred embodiment a metal oxide is also added.



EUROPEAN SEARCH REPORT

Application Number EP 98 10 5081

		RED TO BE RELEVANT	Delevent	01 4001510 4 710 4 07 717
Category	Citation of document with ind of relevant passage	• • • • • • • • • • • • • • • • • • • •	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
D,Y	US 4 797 462 A (M. L 10 January 1989 * column 1, line 7 - * column 5, line 16 * column 10, line 14 * example 2 *	line 12 * - line 21 *	1-9	C08L83/04
Υ .	CHEMICAL ABSTRACTS, 23 September 1991 Columbus, Ohio, US; abstract no. 116011, XP002086757 * abstract * & JP 03 052955 A7 Ma		1-9	·
Y	EP 0 493 887 A (GENE 8 July 1992 * claim 1 *	RAL ELECTRIC)	1-9	
Υ	FR 1 569 201 A (MIDL 30 May 1969 * page 1, line 1 - 1 * page 1, line 15 - * page 6, line 6 - 1 * claims 1,2 * -	ine 3 * line 20 *	1-9	TECHNICAL FIELDS SEARCHED (Int.CI.6) CO8K CO8L
				·
			•	
	The present search report has be	een drawn up for all claims		
	Place of search	Date of completion of the search		Examiner
	BERLIN	4 December 1998	B Hoe	pfner, W
X : part Y : part doci A : tech O : non	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with another ument of the same category anological background i-written disclosure irmediate document	E : earlier patent after the filling D : document cite L : document cite	ciple underlying the document, but publication in the application of for other reasons as same patent familication.	ished on, or

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 98 10 5081

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

04-12-1998

Patent document cited in search report		Publication date	Patent family member(s)		Publication date	
US 47	97462	A	10-01-1989	FR	2603894 A	18-03-1988
				AU	594817 B	15-03-1990
•				AU	7831587 A	17-03-1988
				CA	1285085 A	18-06-1991
			•	DE	3774068 A	28-11-1991
				EP	0264336 A	20-04-1988
				GR	3003427 T	17-02-1993
				JP	1721396 C	24-12-1992
			•	JP	4006745 B	06-02-1992
				JP	63086751 A	18-04-1988
•			•	PT	85676 B	31-05-1990
EP 49	 3887	Α	08-07-1992	CA	2056569 A	18-06-1992
				DE	69117036 D	21-03-1996
				DE	69117036 T	19-09-1996
				ES	2085442 T	01-06-1996
			•	JP	2698268 B	19-01-1998
				JP	4359055 A	11-12-1992
			·	US	5489479 A	06-02-1996
FR 15	69201	Α	30-05-1969	DE	1769537 A	23-12-1970
				GB	1219754 A	20-01-1971